Samples

CRP-containing samples of varying CRP concentration were prepared from a 200 mg/l of recombinant CRP (Fitzgerald) in hCRP depleted serum.

5 Procedure

 $15~\mu l$ of CRP-containing serum diluted 1/20 in test buffer (50 mM borate buffer pH 8.0, 3% BSA, 5% sucrose, 0.15 M NaCl, 0.005% CaCl₂, 0.05% NaN₃) were applied to the application zone of the membrane strip. Then, 15 μl of detection conjugate solution [anti-CRP monoclonal antibody (Fitzgerald) coupled to 0.1 μm TransFluoSpheres-SO4/CHO (633/720 nm) (Molecular Probes Inc.), the above test buffer] were added, the amount of anti-CRP conjugate being 3 μg per test strip which was a 15 x molar excess in relation to the highest standard value. The conjugate addition was followed by a wash with 15 μl of test buffer. The fluorescence of the strip

was then measured. The results are shown in Table 2 below.

Table 2

Peak area obtained
(V x mm)
0.41
0.60
7.51
7.130
8.86
9.42
11.97
10.67
11.70
12.91
14.27
14.16

15

10

The above Examples 1 and 2 thus demonstrate that it is possible to run an assay on undiluted high concentration samples without using huge amounts of reagents when using the methodology of the present invention.